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AMENDMENTS TO THE CLAIMS: This listing of claims will replace prior versions and listings

of claims in the application:

Claims 1-5, 13, 37, 39, 41, 43 and 44 have been amended as follows:

Underlines indicate insertions and strikethrough indicate deletions.

Listing of claims:

1. (Currently Amended) A microporous polymeric article having a target pore

diameter d<sub>v.</sub> comprising an essentially continuous porosity with a controlled void volume controlled frem between 10 to and 90%, wherein pore diameters show are controlled

<u>within a unimodal distribution set to a predefined unimodal peak location corresponding</u>

to a chosen said target pore diameter dy, and wherein a majority of pores has have a

diameter d comprised in a range between d<sub>v</sub>-50% and d<sub>v</sub>+50% within at most ± 50% of

the chosen pore diameter.

2. (Currently Amended) The microporous polymeric article according to claim 1,

wherein the predefined unimodal peak location corresponds to a chosen pore diameter

selected the target pore diameter  $d_v$  is from 20 nm to 500  $\mu m$ .

3. (Currently Amended) The microporous polymeric article according to claim 2,

wherein the predefined unimodal peak location corresponds to a chosen pore diameter

selected the target pore diameter  $d_{\nu}$  is from 1 to 72  $\mu m.$ 

4. (Currently Amended) The microporous polymeric article according to claim 3,

wherein the majority of pores  $\,$  have a diameter  $\underline{d}$  comprised in a range between  $\underline{d}_{v}\text{--}40\%$ 

and  $d_v$ +40% within ± 40% of thechosen pore diameter.

5. (Currently Amended) The microporous polymeric article according to claim 1,

wherein the target pore diameter d<sub>v</sub> is predefined unimodal peak location corresponds to

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a-chosen pore-diameter-selected from 1 to 3  $\mu$ m, and wherein the majority of pores have a diameter <u>d</u> comprised in a range between d<sub>v</sub>-25% and d<sub>v</sub>+25% within ± 25% of the chosen pore-diameter.

- (Withdrawn) The microporous biodegradable polymeric article according to claim 1, wherein the porosity is fully continuous.
- (Withdrawn) The microporous biodegradable polymeric article according to claim 1, wherein the article has a symmetric morphology.
- (Withdrawn) The microporous biodegradable polymeric article according to claim 1, wherein the article has an asymmetric morphology.
- (Withdrawn) The microporous biodegradable polymeric article according to claim
  wherein the article has a closed-cell skin.
- 10. (Withdrawn) The microporous biodegradable polymeric article according to claim 1, wherein at least 95% of said article is made of a biodegradable medical polymer selected from the group consisting of poly(lactic acid), poly(glycolic acid), poly(lactic-coglycolic), polyorthoesters, polycaprolactones, polyanhydrides and their copolymers.
- 11. (Withdrawn) The microporous biodegradable polymeric article according to claim 1, wherein at least 99% of said article is made of a biodegradable medical polymer selected from the group consisting of poly(lactic acid), poly(glycolic acid), poly(lactic-coglycolic), polyorthoesters, polycaprolactones, polyanhydrides and their copolymers.
- 12. (Withdrawn) The microporous biodegradable polymeric article according to claim 1, wherein said article is essentially made of a biocompatible, implantable polymer.
- 13. (Currently Amended) A microporous polymeric article <u>having a target pore diameter d<sub>v</sub></u> comprising an essentially continuous porosity with a controlled void volume from 10 to 90%, wherein pore diameters show a unimodal distribution set at a predefined unimodal peak location corresponding to a-chosen said target pore diameter d<sub>v</sub>, and wherein a majority of pores have[] a diameter d within at most ± 50% of at least

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d<sub>v</sub>-50% and at most d<sub>v</sub>+50% of the chosen pore diameter, prepared according to a method comprising the steps of:

a) determining a target pore diameter d<sub>v</sub>:

a) b) selecting at least one polymer  $A_{\bar{\tau}}$  and at least one polymer  $B_{\bar{\tau}}$  at least

partially immiscible with A, according the target pore diameter d<sub>v</sub> determined in

step a);

b) c) melt blending the selected polymers from step a) b), thereby preparing a

polymer blend, wherein said polymers A and B have an essentially continuous

morphology;

 $\underline{\text{e)-}\underline{\text{d)}}}$  cooling said polymer blend to room temperature, thereby retaining its

morphology; and (a) extracting said polymer B, at least partially, from the polymer blend by dissolving said polymer B in a solvent that is a non-solvent of

polymer A.

14. (Withdrawn) A method of preparation of a microporous biodegradable polymeric

article, comprising the steps:

a) selecting at least one biodegradable polymer A, one polymer B,

biodegradable or not, at least partially immiscible with A, and a polymeric

compatibilizer C for A and B;

b) melt blending the selected polymers from step a) and the compatibilizer C,

thereby preparing a compatibilized polymer blend, wherein said polymers A and

B have an essentially continuous morphology;

c) cooling said polymer blend to room temperature, thereby retaining its

morphology; and

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d) extracting said polymer B and said compatibilizer C, at least partially, from the polymer blend by dissolving them in a solvent that is a non-solvent of polymer

A.

wherein said polymeric article has an essentially continuous porosity with a void volume from 10 to 90%, wherein pore diameters show a unimodal distribution set to a

predefined unimodal peak location corresponding to a chosen pore diameter, and

wherein a majority of pore has a diameter within  $\pm\,50\%$  of the chosen pore diameter.

15. (Withdrawn) The method according to claim 14, wherein said polymer A is a

biodegradable medical polymer.

16. (Withdrawn) The method according to claim 15, wherein said polymer A is an

aliphatic polyester.

17. (Withdrawn) The method according to claim 15, wherein said polymer A is selected from the group consisting of poly(lactic acid), poly(glycolic acid), poly(lactic-co-

glycolic), poly(hydroxyalkanoates), polyorthoesters, polycaprolactones, polydioxanone,

polyanhydrides and their copolymers.

18. (Withdrawn) The method according to claim 14, wherein said polymer B is a non-

biodegradable polymer.

19. (Withdrawn) The method according to claim 14, wherein said polymer B is a

biodegradable medical polymer.

20. (Withdrawn) The method according to claim 19, wherein said polymer B is

selected from a group consisting of poly(lactic acid), poly(glycolic acid), poly(lactic-co-

 $\hbox{glycolic), poly(hydroxyalkanoates), polyorthoesters, polycaprolactones, polyanhydrides}\\$ 

and their copolymers.

21. (Withdrawn) The method according to claim 14, wherein said compatibilizer C is

a polymeric compatibilizer.

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22. (Withdrawn) The method according to claim 21, wherein said compatibilizer C is a copolymer of A and B.

 (Withdrawn) The method according to claim 14, wherein said polymers A and B are fully immiscible.

24. (Withdrawn) The method according to claim 14, wherein said polymer blend is co-continuous at more than 90%.

 (Withdrawn) The method according to claim 14, wherein said polymer blend may contain one or more additives.

26. (Withdrawn) The method according to claim 14, wherein said polymer blend is submitted to a further step of controlled annealing between steps b) and c), thereby increasing the pore size of the porous article.

27. (Withdrawn) The method according to claim 14, wherein said polymer blend is submitted to controlled cooling rates in step c).

28. (Withdrawn) The method according to claim 14, wherein said polymer blend is further shaped into a geometrical form between steps b) and c).

29. (Withdrawn) The method according to claim 28, wherein said polymer blend is further shaped in a mold or die, between steps b) and c).

30. (Withdrawn) The method according to claim 28, wherein said polymer blend is shaped by injection molding, between steps b) and c).

31. (Withdrawn) The method according to claim 28, wherein said polymer blend is formed by extrusion, between steps b) and c).

32. (Withdrawn) The method according to claim 28, wherein said polymer blend is formed by melt spinning between steps b) and c).

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(Withdrawn) The method according to claim 14, wherein said polymer blend is 33. submitted to a mechanical stress that orients the porosity in at least one specific

direction, between steps b) and c).

34. (Withdrawn) The method according to claim 14, wherein said polymer blend is

submitted to a mechanical stress that orients the porosity in at least one specific

direction, during step c).

35 (Withdrawn) The method according to claim 14, wherein said polymeric article is

further submitted to a controlled immersion in a solvent for its polymer A after step d),

thereby creating a closed-cell skin.

36. (Withdrawn) The method according to claim 14, wherein said polymer blend is

further submitted to a controlled immersion in a common solvent for A and B between steps c) and d), thereby creating an asymmetric open-cell morphology in the porous

article.

37.

(Previously presented) A method of tissue engineering utilizing a microporous

article according to any one of claims 1 to 5 and 13.

38 (Withdrawn) The use of a microporous biodegradable article obtained by the

method according to any of claims 14-36 in tissue engineering.

39. (Previously presented) A method of controlled release utilizing a microporous

article according to any one of claims 1 to 5 and 13 as a substrate.

40 (Withdrawn) The use of a microporous biodegradable article obtained by the

method according to any of claims 14-36 as a substrate for controlled release

applications.

41. (Previously presented) A method of forming an implantable medical device

utilizing a microporous article according to any one of claims 1 to 5 and 13.

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42. (Withdrawn) The use of a microporous biodegradable article obtained by the method according to any of claims 14-36 as an implantable medical device.

- (Previously presented) The microporous polymeric article according to any one of claims 1 to 5, said article being biodegradable.
- 44. (Currently amended) The microporous polymeric article according to claim 13, wherein step a) b) further comprises selecting a polymeric compatibilizer C for A and B; step c) b) comprises melt blending the selected polymers from step a) and the compatibilizer C, thereby preparing a compatibilized polymer blend; and step e) d) comprises extracting said polymer B and said compatibilizer C, at least partially, from the polymer blend by dissolving them in a solvent that is a non-solvent of polymer A.